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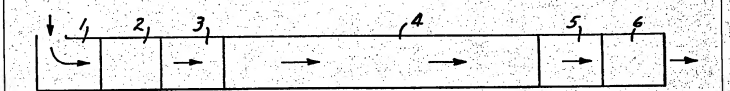
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(54) Title: GRAFT COPOLYMERS CONTAINING FLUOROALIPHATIC GROUPS



(57) Abstract

A fluorochemical graft copolymer comprising: a base polymer comprising polymerized units derived from monomers having terminal lefinic double bonds, having a moiety comprising a fluoroaliphatic group grafted thereto. Also disclosed are processes for preparing such fluorochemical graft copolymers, forming and annealing methods for enhancing the surface activity of such a graft copolymer, polymer blends comrpising such a fluorochemical graft copolymer and a matrix polymer that is miscible with the base polymer, and a method for reducing the surface energy of a polymer comprising polymerized units derived from monomers having terminal olefinic double bonds.

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GRAFT COPOLYMERS CONTAINING FLUOROALIPHATIC GROUPS

BACKGROUND OF THE INVENTION

Technical Field

This invention relates to continuous processes using wiped-surface reactors for free radical 10 graft polymerization of polyolefins, and to graft copolymers thereby produced. In another aspect, this invention relates to polymerizable olefins containing fluoroaliphatic groups, and copolymers thereof.

15 Description of the Related Art

Processing or production of polymeric resins using wiped-surface reactors such as screw extruders and twin screw extruders is well known (such processing is often called reactive extrusion). Twin screw 20 extruders and their use in continuous processes such as graft polymerization, alloying, bulk polymerization of vinyl monomers, and condensation and addition reactions are generally described in Plastics Compounding Jan./Feb. 1986, pp. 44-53 (Eise et al.) and Plastics 25 Compounding, Sept./Oct. 1986, pp. 24-39 (Frund et al.). Graft reactions are said to be carried out by first melting a polymeric species in the initial stages of an extruder, injecting a peroxide catalyst into the extruder, and mixing in a monomer under high shear 30 conditions. Advantages of the twin screw extrusion process are said to include narrow distribution of molecular weight, improved melt-flow properties, consistent process control, and continuous processing.

Graft polymerization reactions of

35 polyolefins with various monomers using wiped-surface reactors are known. Such grafting is said to be useful in providing a polymer adduct with functionality to allow further modification of structure and properties. General mechanistic proposals regarding formation of

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these "mechanochemically synthesized" adducts are discussed in connection with grafting of maleic anhydride onto polypropylene in Polymer Prep., 1986, 27, 89 (Al-Malaika).

A number of particular free radical graft polymerization reactions have been reported. For example, U.S. Pat. No. 3,177,270 (Jones et al.) discloses a process for preparing graft copolymers by mixing an olefin polymer at a temperature between 110°C 10 and 250°C while contacting the polymer with a minor proportion of a mixture comprising a monovinyl aromatic compound and optionally one or more other monomers such as acrylic acid, methacrylic acid, acrylonitrile, methyl methacrylate, methacrylonitrile, or maleic 15 anhydride, the mixture having dissolved therein an organic peroxide.

British Pat. No. 1,292,693 (Steinkamp et al.) discloses use of a single-screw extruder to graft monomers such as maleic anhydride and acrylic acid onto 20 polyolefins such as polypropylene in the presence of a suitable free radical initiator such as an organic peroxide. The product graft copolymers are said to have a melt flow rate (MFR) of at least 50% greater than the MFR of the base polymer.

U.S. Pat. No. 4,003,874 (Ide et al.) discloses modified polyolefins obtained by adding an unsaturated carboxylic acid or an anhydride thereof and an organic peroxide to a polyolefin and melting these components in an extruder. The polyolefin so obtained 30 is said to adhere to glass fibers.

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U.S. Pat. No. 4,146,529 (Yamamoto et al.) discloses a process for production of modified polyolefins by combining a polyolefin with one or more carboxylic acids or their anhydrides in an extruder in 35 the presence of a radical producing agent and an organosilane.

U.S. Pat No. 4,228,255 (Fujimoto et al.) discloses a meth d for crosslinking a polyolefin, the polyolefin being a low density polyethylene or a

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polyolefin mixture containing a low density polyethylene, comprising reacting the polyolefin with an organic silane and an organic free radical initiator to form a silane-grafted polyolefin, then mixing the silane-grafted polyolefin with a silanol condensation catalyst. The mixture is extruded with heating in a single-screw extruder to provide a crosslinked polyethylene.

10 <u>SUMMARY OF THE INVENTION</u>

This invention provides a fluorochemical graft copolymer comprising: a base polymer comprising polymerized units derived from monomers having terminal olefinic double bonds, having a moiety comprising a fluoroaliphatic group grafted thereto. The grafted fluoroaliphatic group is generally derived from a fluorochemical olefin comprising a fluoroaliphatic group and a polymerizable double bond.

The fluoroaliphatic group of the
20 fluorochemical olefin is generally bonded to the
polymerizable double bond through a linking group.
Such fluorochemical olefins can be represented by
Formula I below:

 $(R_f)_{\bullet}Q(CR=CH_2)_{\bullet}$

30

wherein R is hydrogen, trifluoromethyl, or straight chain or branched chain alkyl containing 1 to about 4 carbon atoms;

a is an integer from 1 to about 10;
b is an integer from 1 to about 6;
Q is an (a+b)-valent linking group that does not substantially interfere with free radical polymerization; and

35 R_f is a fluoroaliphatic group comprising a fully fluorinated terminal group containing at least seven fluorine atoms.

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This invention also provides a process for preparing the fluorochemical graft copolymers described above, which process comprises the steps of:

- (1) feeding to a reactor materials 5 comprising:
 - (a) a base polymer comprising polymerized units derived from monomers having terminal olefinic double bonds;
 - (b) an effective amount of a free radical initiator system comprising one or more free radical initiators; and
 - (c) a fluorochemical olefin as described above, wherein all materials are substantially free of oxygen;
 - (2) reacting the materials in the reactor to provide a graft copolymer; and
 - (3) withdrawing the graft copolymer from the reactor.

Preferably, the base polymer is fed to the 20 reactor in a region of the reactor preceding or coincident with the region in which the initiator system is fed and the fluorochemical olefin is fed to the reactor in a region of the reactor subsequent to the region in which the initiator is fed.

A preferred embodiment of the process of the invention involves use of a least two free radical initiators to maximize the number of grafted fluorochemical moieties. Desired thermoplastic, melt-flow, and low surface energy properties of the 30 resultant graft copolymer can thus be optimized.

In another aspect, this invention provides methods of enhancing the surface activity of a fluorochemical graft copolymer film. One such method is an annealing method comprising the steps of:

- a) providing a surface comprising a fluorochemical graft copolymer as described above, and
- annealing the surface by heating it at a temperature and for a time effective to increase the amount of the grafted fluorochemical at the surface.

This invention also provides a forming method for controlling the amount of grafted fluorochemical at the surface of a composition comprising a graft copolymer of the invention, 5 comprising the steps of:

- (1) selecting a surface that is made of a material that will control the amount of grafted fluorochemical at the surface of the composition; and
- (2) forming the composition against the
 10 surface selected in step (1).

This invention also provides polymer blends comprising a fluorochemical graft copolymer as described above and a matrix polymer that is miscible with the base polymer of the graft copolymer.

- Further, this invention provides a method for reducing the surface energy of a polymer, comprising the step of grafting to said polymer a fluorochemical olefin comprising a fluoroaliphatic group and a polymerizable double bond.
- Relatively little fluorochemical olefin is required for the preparation of the graft copolymers of the invention, and the forming and annealing methods further reduce the amount of fluorochemical needed for a particular application.
- Graft copolymers of the invention have lower surface energies than the corresponding base polymers, and they exhibit desirable thermoplastic, melt flow, and release properties. Moreover, these graft copolymers are useful in applications where oil repellency and solvent resistance are desirable.

BRIEF DESCRIPTION OF THE DRAWING

The Drawing is represented by FIGS. 1-3.

FIG. 1 is an exemplary flow diagram of the process of the invention. Ancillary equipment such as pumps and valves, has not been illustrated, and secondary process streams such as utility lines (e.g., cooling water) have be n omitted.

FIG. 2 is a flow diagram of a counter-rotating twin screw extruder useful in this invention.

FIG. 3 is a flow diagram of another counter-5 rotating twin screw extruder useful in the process of this invention.

DETAILED DESCRIPTION OF THE INVENTION

A fluorochemical graft copolymer of the invention comprises a base polymer having a moiety comprising a fluoroaliphatic group grafted thereto. Generally a plurality of the grafted moiety is present in a graft copolymer of the invention.

The fluoroaliphatic group is generally

15 derived from a fluorochemical olefin comprising a
fluoroaliphatic group and a polymerizable double bond.

The grafting occurs through the polymerizable double
bond. The double bond is of course not present in the
product graft copolymer of the invention; rather, in

20 the grafting process the double bond becomes a
saturated link between the base polymer and the
fluoroaliphatic group. In the instant specification
and claims a reference to a fluoroaliphatic group
grafted through a double bond designates the presence

25 of such a saturated link and does not designate the
presence of olefinic unsaturation in the grafted moiety
as it is incorporated in the graft copolymer.

Suitable base polymers include polymers comprising polymerized units derived from monomers having terminal olefinic double bonds. This class of polymers is known to those skilled in the art and includes polymers such as polymethyl methacrylate, poly-4-methylpentene, polypropylene, polybutylene, polystyrene, polyethylene, polybutadiene, and copolymers such as ethylene/vinyl acetate copolymer and thylene/butyl acrylate copolymer, and the like, and mixtures and blends thereof. Such polymers of any molecular weight are suitable. Polymers with a wide range of melt index values (e.g., from about 0.1 to

about 500) are suitabl. Melt index values are determined by the American Society for Testing Materials method ASTM D-1238. Due to their relatively low viscosity and ability to diffuse to the surface of a polymer blend, low molecular weight polyolefins, preferably with melt indices of at least about 20, (e.g., low molecular weight polyethylene and polypropylene), are particularly useful as base polymers for graft copolymers of the invention that are intended for use in the polymer blends of the invention (described in detail below).

Fluorochemicals that can be grafted to the base polymer include known fluorochemical olefins that comprise a fluoroaliphatic group and a polymerizable 15 double bond. Fluorochemical olefins suitable for use in the invention include those disclosed in, for example, U.S. Pat. Nos. 2,642,416 (Ahlbrecht et al.), 2,803,615 (Ahlbrecht et al.), 2,841,573 (Ahlbrecht et al.), 3,102,103 (Ahlbrecht et al.), 3,282,905 (Fasick 20 et al.), 3,304,278 (Hauptschein et al.), 3,378,609 (Fasick et al.), 3,384,627 (Anello et al.), 3,386,977 (Kleiner), 3,392,046 (Marder), 3,407,183 (Farah et al.), 3,514,420 (Katsushima et al.), 3,532,659 (Hager et al.), 3,544,663 (Hauptschein et al.), 3,546,187 25 (Tandy), 3,547,861 (Anello et al.), and 3,578,487 (Knell et al.). The list above is intended to be merely exemplary and not exhaustive of patents disclosing suitable fluorochemical olefins. Generally, suitable fluorochemical olefins comprise a 30 fluoroaliphatic group bonded through a linking group to a polymerizable double bond and can be represented by the general Formula I below:

 $(R_f)_{a}Q(CR=CH_2)_{b}$

I

wherein a, b, R, R, and Q are as defined above.

In a compound of Formula I, a and b are integers representing the number of fluoroaliphatic

groups and the number of olefinic groups, respectively, in the fluorochemical olefin. The value of a can be 1 to about 10, preferably 1 to about 6, more preferably 1 to about 3, and most preferably 1. The value of b can be 1 to about 6, preferably 1 to about 3, and more preferably 1. R in a compound of Formula I is hydrogen, trifluoromethyl, or lower alkyl (i.e., straight chain or branched chain alkyl of 1 to about 4 carbon atoms).

Q is an (a+b)-valent organic moiety that can have a wide variety of structures, for example, alkylene, e.g., methylene, ethylene, cyclohexylene, arylene, e.g., phenylene, and combinations thereof, e.g. xylylene, or combination of such moieties with 15 such heteroatom-containing moieties as oxy, thio, aza, carbonyl, sulfonyl, sulfoxy, sulfonamido, carboxamido, urylene, carbamato, and imino, and combinations thereof such as sulfonamidoalkylene, carboxamidoalkylene, oxydialkylene, alkylenecarbamato and the like. The 20 particular structure of Q for a particular fluorochemical olefin is not unduly critical to this invention. Q can therefore be selected by virtue of ease of preparation and, for example, commercial availability of the fluorochemical olefin or the 25 particular reactants used in preparing the fluorochemical olefin.

R_f is a fluoroaliphatic group that is a fluorinated, stable, inert, non-polar, preferably saturated, and both hydrophobic and oleophobic. R_f can be straight chain, branched chain, or, if sufficiently large, cyclic, or a combination thereof, such as alkylcycloalkyl. The fluoroaliphatic group can also include catenary oxygen, sulfur, or nitrogen.

Generally R_f will have 3 to about 20 carbons atoms,

preferably 6 to about 12 carbon atoms, and will contain about 40 to about 78 weight percent, preferably about 50 to about 78 weight percent, carbon-bound fluorine. The terminal portion of the R_f group is fully fluorinated and contains at least 7 fluorine atoms.

Exemplary terminal portions include $-CF_2CF_2CF_3$, $-CF(CF_3)_2$, $-CF_2SF_5$, and the like. Preferr d R_f groups are fully or substantially fully fluorinated, as in the case where R_f is perfluoroalkyl (i.e., C_nF_{2n+1}).

Several particular exemplary compounds of Formula I are shown below:

 $C_8F_{17}SO_2N (C_4H_9) CH_2CH_2O (O) CCH=CH_2$ $C_8F_{17}SO_2N (C_2H_5) CH_2CH_2O (O) CCH=CH_2$ $C_8F_{17}SO_2N (C_2H_5) CH_2CH_2O (O) CC (CH_3)=CH_2$

10 $C_7F_{15}CH_2O(O)CCH=CH_2$ $C_7F_{15}CH_2O(O)CC(CH_3)=CH_2$ $C_8F_{17}SO_2N(C_2H_5)CH_2CH=CH_7$

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 $C_8F_{17}SO_2N$ (CH₃) CH₂CH₂O (O) C (CH₃) =CH₂

 $C_8F_{17}SO_2N(CH_3)CH_2CH_2CH_2CH_2O(O)C(CH_3)=CH_2$

15 $C_8F_{17}SO_2N (CH_3) CH_2CH=CH_2$ $C_8F_{17}SO_2N (CH_2) CH_2CH_2OCH=CH_2$ $C_4F_9OC_3F_6OCF (CF_3) CH_2O (O) CCH=CH_2$ $C_8F_{17}C_2H_4SO_2N (CH_3) C_2H_4O (O) CCH=CH_2$

 $C_8F_{17}(CH_2)_{10}O(O)CCH=CH_2$ $C_8F_{17}CH_2CH_2O(O)CCH=CH_2$ $C_6F_{13}C_2H_4SC_2H_4O(O)CCH=CH_2$

 $(C_3F_7)_3CCH_2O(O)CCH=CH_2$

 $(C_4F_9CO)_2NCH_2CH_2O(O)CC(CH_3)=CH_2$

CF₂-CF₂

CF₂-CF₂

NCF₂CF₂C(0) N(CH₃) CH₂CH₂O(0) CCH=CH₂

CF₂-CF₂

 $SF_5(CF_2)_4CH_2OC(O)CH=CH_2$

35 $F_{3}C-C-CF_{3}$ CF_{3} $CH_{2}=CHC(O)OCCH_{2}CH_{2}CH_{2}N(CH_{3})SO_{2}-C-F$ CF_{3}

In a preferred embodiment, the graft copolymer comprises a polymeric backbone comprising polymerized units derived from monomers having terminal olefinic double bonds, having bonded thereto a moiety of the formula

-CH₂CHR-Q-Rf

wherein Q is a divalent linking group that does not substantially interfere with free radical polymerization, and R and R are as defined above.

10 Preferably a graft copolymer of the invention comprises about 0.1% to about 20%, more preferably 0.5 to about 10% by weight of grafted fluorochemical olefin. In a process of the invention as described below it is preferred to use like 15 quantities of fluorochemical olefin, i.e., about 0.1 to about 20% or more by weight, more preferably 0.5 to about 10% weight based on the weight of the base polymer.

In a process of the invention, the base
20 polymer and a fluorochemical olefin are reacted in the
presence of an initiator system comprising one or more
free radical initiators to provide a fluorochemical
graft copolymer. The initiator system serves to
initiate free radical grafting of the fluorochemical
25 olefin onto the base polymer.

Many initiators are known. Suitable initiators include: hydroperoxides such as cumene, t-butyl, and t-amyl hydroperoxides, and 2,5-dihydroperoxy-2,5-dimethylhexane; dialkyl peroxides such as di-t-butyl, dicumyl, and t-butyl cumyl peroxides, 2,5-dimethyl-2,5-di(t-butylperoxy)hexane, and 2,5-dimethyl-2,5-di(t-butylperoxy)hex-3-yne; peroxyesters such as t-butyl perbenzoate and di-t-butyl-diperoxy phthalate, diacyl peroxides such as benzoyl peroxide and lauroyl peroxide; peroxyketals such as n-butyl-4,4-bis(t-butylperoxy)valerate and 1,1,-di-t-butylperoxy-3,3,5-trimethylcyclohexane; and azo compounds such as azoisobutyronitrile.

Reaction conditions under which a graft copolymer of the invention can be prepared typically involve heating at about 150°C to about 250°C.

Reactants typically have a residence time of about 1 to 5 20 min. It is therefore difficult to select a single initiator with a decomposition rate such that initiating radicals will be present in a substantial concentration for a prolonged period of time when a low concentration of initiator is used. It is therefore 10 preferred to use a mixture of at least two initiators as an initiator system. Proper selection of the components of the initiator system overcomes the above-discussed difficulty with single initiators, and allows control and optimization of the physical properties of the product graft copolymer.

Generally it is preferred that each initiator in an initiator system have a rate of decomposition substantially different from those of the other initiators in the initiator system. For example, in a process with a residence time of about 5-10 minutes at a temperature of about 200°C, an initiator system wherein one initiator has a half-life of about 30 seconds and the other initiator has a half-life of about 2 minutes has been found suitable.

25 Preferred initiator systems include mixtures comprising from about 40% to about 60% by weight of 2,5-dimethyl-2,5-di(t-butylperoxy) hexane, (such as that commercially available as LUPERSOLTM 101 from Pennwalt Corporation) and from about 60% to about 40% 30 by weight of an initiator such as 2,5-dimethyl-2,5-di(t-butylperoxy) (such as that commercially available as LUPERSOLTM 130, Pennwalt Corporation), t-butylhydroperoxide, or di-t-butylperoxide. Initiator decomposition rates are 35 temperature dependent, and other particular initiator systems and preferred concentrations thereof can be selected by those skilled in the art consistent with the temperature of the reaction and the residence time of the reactants.

Total initiator concentration in a process of the invention is preferably from about 0.1% to about 1%, more preferably from about 0.25% to about 0.5% based on the weight of the base polymer.

Fluorochemical graft copolymers of the invention can be prepared using various well known reactors such as stirred tank reactors, tubular reactors, and extruders. Graft copolymers are preferably made by a process involving a wiped-surface 10 reactor. A wiped surface reactor comprises a shell or vessel that contains at least one rotor having a wiping portion located close to the inside surface of the shell and a root portion spaced further from the shell than the wiping portion. As the rotor is rotated, the 15 wiping portion passes close enough to the inside surface of the shell to clean the surface and form a seal when the reactor contains monomer and/or polymer but not so close as to cause permanent deformation of either the rotor or shell. It is necessary that the 20 root surface of the rotor also be wiped or cleaned continuously during the operation of the reactor.

Intermeshing twin screw extruders can be used as wiped surface reactors. The screws function as the rotor and the flight lands function as the wiping 25 portion, while the screw root surface between the flight lands functions as the root surface. Clearances between the inside of the barrel wall of the extruder and the flight lands of the screws are preferably from about 0.25 to 0.5 mm. Although co-rotating twin screw 30 extruders can be used, counter-rotating twin screw extruders are preferred. The counter-rotating extruder acts as a positive displacement pump conveying the reactant stream, and it also behaves as a series of small mixing zones or continuous stirred tank reactors. The counter-rotating twin screw extruder also gives good control over melting, mixing, and reaction temperatures.

Preferably, screws of a counter-rotating twin screw extruder are divided into segments, i.e.,

xtruder screws can be composed of a number of s parate screw segments that fit onto a common drive shaft by a keyway and can be disassembled and rearranged in various orders and configurations. It is also possible to use screw segments having multiple (e.g., two or three) starts and various pitch, and one or more screw segments can be reversed to increase mixing. Residence time of the reactants, and properties of the resultant product, can therefore be varied by selection of screw pitch and/or screw speed (i.e., screw rpm).

Furthermore, each particular zone of a twin screw extruder can be independently heated or cooled by external heating or cooling means, allowing further control of reaction conditions.

Use of a wiped surface reactor in the 15 invention is discussed referring to FIG. 1. A base polymer can be fed in a region of the reactor coincident with the region in which the initiator system is fed. For example, the desired base polymer, 20 preferably in pellet form, can be wetted with a free radical initiator system and purged with an inert gas such as nitrogen, helium, argon or the like, to render the material substantially free of oxygen (i.e., oxygen, if present, is present in an amount such that it does not significantly affect the desired free radical polymerization reactions). This material can be fed at a predetermined rate into feed zone 1 of the wiped surface reactor. It is preferred, however, to feed the base polymer in a region of the reactor 30 preceding the region in which the initiator system is fed. Feed zone 1 typically comprises a feed throat, into which base polymer can be fed into the upstream end, and into which the initiator system can be fed at the downstream end.

Dase polymer and the initiator involves use of a 2-component feed zone consisting of a base polymer feed zone into which base polymer is fed, followed in sequence by a separate initiator feed zone into which

the initiator is fed. The extruder is preferably starve fed, i.e., all material fed into the feed zone is conveyed into initiator/melt zone 2 of the extruder, and nothing is held up in feed zone 1.

Feed rates can vary with the size of the reactor and for any given size of reactor, one skilled in the art will be able to determine suitable feed rates. As an example, when a 34 mm counter-rotating twin screw extruder is used, feed rates are preferably from about 0.4 Kg/h to about 9 Kg/h. The feed zone screw preferably has a high pitch (e.g., 20 mm) to accommodate base polymer pellets. The feed zone can, if desired, be operated in a temperature controlled manner, depending on the reactants, reaction conditions and the like. Generally, it is suitable to maintain the feed zone of the extruder in a temperature range from about 10°C to about 50°C, depending on the base polymer used.

In initiation/melt zone 2, the initiator

20 system and base polymer are mixed and heated to
initiate radical chain reactions. Preferred
temperatures will depend on the particular base polymer
and initiator system, but generally temperatures in the
range between 150°C and about 250°C are suitable.

In monomer addition zone 3, a
nitrogen-purged fluorochemical olefin is added, usually
by a high pressure pump and under an inert atmosphere.
The fluorochemical olefin is generally fed as a liquid
or as a solution in an inert solvent (e.g., decane,
30 toluene, tetrahydrofuran or the like. Preferred feed
rates are variable, and when a LEISTRITZTM 34 mm
counter-rotating twin screw extruder is used, feed rate
is preferably about 4 g/h to about 180 g/h. It is
preferred to maintain the monomer addition zone at a
35 temperature of about 150°C to about 250°C.

Grafting proceeds in reaction zone 4. The reaction zone is heated. As with the initiator/melt zone, the preferred temperature will depend on the particular base polymer and initiator system used.

Further, the pr f rr d temperature of the reaction zone will depend on the particular base polymer and initiator system used and on the intended residence time in the reaction zone. Generally, temperature in the range of 150°C to 250°C and residence times in the range of 1 minute to about 10 minutes are suitable.

In reactions where residual solvent or monomer remains, it is preferred to remove it from the grafted product by venting. This can be done in devolatilization zone 5, where a vacuum (e.g., about 10 kPa absolute pressure) can be applied to a vent line. The resultant product is then passed through block zone 6, which conveys the product graft copolymer for any further processing as desired, e.g., shaping in a die, 15 quenching in a suitable quenching liquid, or pelletizing for convenience of handling and/or storage.

The graft copolymers of the invention can find application where good release properties, oil repellency, water repellency, solvent resistance, and other properties of fluorochemicals are advantageous.

The amount of grafted fluorochemical at the surface (i.e., the surface fluorochemical density) of a composition comprising a graft copolymer of the invention can be increased by the annealing method of 25 the invention. In such a method, a graft copolymer or a film thereof is annealed at a temperature and for a time effective to increase the amount of grafted fluorochemical at the surface. Effective temperature and time will bear an inverse relationship to one 30 another and a wide range of conditions will be suitable. Generally, annealing at a temperature between about 50°C and about 160°C for a period of several minutes to several hours provides a composition with an increased amount of grafted fluorochemical at 35 the surface. Annealing apparently allows the fluorochemical in the graft copolymer to migrate to the surface with a resultant increase in surface activity, improved releas properties, and improved oil and water repellency and solvent resistance.

The annealing method, by increasing surface fluorochemical density, can also serve to minimize the amount of grafted fluorochemical olefin that must be present in a graft copolymer of the invention for a particular application.

Another method of controlling the surface fluorochemical density of a composition comprising a graft copolymer of the invention is a forming method, wherein the composition is formed (e.g., as during 10 molding or during pressing into a film) against a selected forming surface. The effect of the selected forming surface on surface fluorochemical density is as follows: Generally, surface fluorochemical density is greater when a composition is formed against a 15 fluorochemical surface, e.g., polytetrafluoroethylene (PTFE, TEFLON™ polytetrafluoroethylene, DuPont) or a surface made of a graft copolymer of the invention, than when it is formed against a polyimide surface (e.g., KAPTONTM polyimide, DuPont). Forming against a 20 chrome surface results in a lower surface fluorochemical density than forming against either polytetrafluoroethylene or polyimide. Through the use of a forming surface having several distinct regions, each of an independently selected material, the forming 25 method of the invention can be used to provide a surface with regionally controlled surface fluorochemical density and hence regionally controlled release properties, oil repellency, and solvent resistance. Further, since the forming method can 30 increase surface fluorochemical density, it can be used to minimize the amount of grafted fluorochemical olefin that must be present in a graft copolymer of the invention for a particular application.

A graft copolymer of the invention can be

35 blended with a matrix polymer that is miscible with the
base polymer of the graft copolymer in order to form a
polymer blend. The base polymer itself is of course a
suitable matrix polymer for use in a polymer blend of
the invention. Furthermore, compilations of polymer

miscibility data are commonly available. Therefore, suitable matrix polymers, i.e., those that are miscible with the particular base polymer, can be easily selected by those skilled in the art. In a polymer 5 blend of the invention the fluorochemical graft copolymer serves as a fluorochemical additive that imparts improved release properties to the matrix polymer. Also, the physical properties (e.g., viscosity, impact strength) of the graft copolymer can be improved by the matrix polymer. Further, the tendency of the graft copolymer to become "physically crosslinked" with the miscible matrix polymer causes the properties imparted by the fluoroaliphatic groups to be more durable than those imparted by a monomeric fluorochemical additive.

A polymer blend of the invention comprises a fluorochemical graft copolymer of the invention in an amount sufficient to impart to the blend the release properties of the fluorochemical graft copolymer. The 20 amount of fluorochemical graft copolymer that constitutes a sufficient amount will vary with the fluorine content of the graft copolymer, and the preferred amount will vary according to the intended use of the polymer blend, Generally, a polymer blend of the invention preferably comprises at least about 1 percent, more preferably at least about 10 percent, and most preferably at least about 30 percent by weight of graft copolymer based on the weight of the matrix polymer.

Objects, features, and advantages of this invention are further illustrated by the following examples. The particular materials and amounts thereof recited in these examples, as well as other conditions and details, should not be construed to unduly limit this invention.

In the following xamples all parts and percentages are by weight unless otherwise specified and all t mperatures are in degr es Celsius unless otherwise indicated.

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EXAMPLES

Examples 1-9

Polypropylene resin base polymer (DYPROTM 8771 pellets, melt index: 9, commercially available from Fina Co., Houston, TX) was mixed in a 5-gallon shaker with 0.25% by weight of 90% liquid 2,5-dimethyl-2,5-di(t-butylperoxy)hexane (LUPERSOLTM 101, Pennwalt Corporation, Philadelphia, PA) and 0.25% by weight of one of the following: (a) 90-95% 2,5-dimethyl-2,5-di(t-butylperoxy)hex-3-yne (LUPERSOLTM 130, Pennwalt); (b) 90% liquid 5-butyl hydroperoxide (LUPERSOLTM 90, Pennwalt); (c) 98.5% liquid di-t-butyl peroxide (Pennwalt).

The base polymer/initiator mixture was then

15 purged with nitrogen for 30 minutes. The resin in the
feed hopper and in the feed zone of the extruder was
kept under nitrogen purge and the base
polymer/initiator mixture was fed with an augered feed
means into the feed zone of a 34 mm counter-otating

20 LEISTRITZTM (LEISTRITZ LSM 30.34 GG, Nurenburg, Germany)
twin-screw extruder (length to diameter, L/D = 35:1)
configured as described below with reference to FIG. 2.

fig. 2 shows a twin-screw extruder with a feed hopper 10, feed zone 12, and a heated barrel that comprises: an initiation/melt zone comprising barrel section 14; a reaction zone comprising a monomer feed zone (barrel section 16) and barrel sections 18, 20, 22, 24, and 26; a devolatilization zone comprising barrel section 28; and a block zone comprising barrel sections 30 and 32. Each barrel section is 120 mm long, and the extruder has a total length of 1200 mm.

Transducer ports (e.g., T4 represents transducer number 4 located in barrel section 24) are located at 30 mm, and/or 90 mm into each heated barrel section. Thermocouple ports are located at 60 mm into each heated barrel section. Melt temperatures of 180°C, 200°C, and 220°C were used with each initiator system. Polymer/initiator flow rate was kept at about 40-50 g/min. N-butyl perfluorooctanesulfonamidoethyl

acrylate (Bufosea) was purged with nitrogen and added to a nitrogen-purged displacement pump, and added in heated barrel section 16, 270 mm from the start of the screws, at a rate of 1 mL/min. Heated barrel section 28 was vented under vacuum. Product graft copolymers were conveyed from the block zone (barrel sections 30 and 32), which was maintained at 180°C, into a water bath and fed into a pelletizer to afford generally cylindrical beads of 3 to 4 mm in length and a diameter of about 1 mm. Reaction conditions are summarized in TABLE 1, wherein L101 designates LUPERSOLTM 101 initiator, L130 designates LUPERSOLTM 130 initiator, TBHP designates t-butyl-hydroperoxide, and DTBP designates di-t-butyl peroxide.

15

TABLE 1

<u>Example</u>	Melt Temp(°)	Initiator <u>Mixture</u>	Polymer Flow Rate (g/min)
20 1	180	L101/L130	48.7
2	200	L101/L130	45.0
3	220	L101/L130	41.4
4	180	L101/TBHP	46.4
5	200	L101/TBHP	44.4
25 6	220	L101/TBHP	44.0
7	180	L101/DTBP	44.0
8	200	L101/DTBP	43.0
9	220	L101/DTBP	35.0

30 Grafting was confirmed by two methods. In the first method, samples in TABLE 1 were extracted in boiling tetrahydrofuran (THF) in a conventional extraction apparatus until a stable weight loss was reached. The polymer was analyzed by infrared spectroscopy before and after extraction. Grafting was confirmed by the presence of strong absorption band at 1720 cm⁻¹ after extraction, which corresponds to the carbonyl absorbance of the acrylate moiety.

Samples of the graft copolymers with 40 thickness of about 0.13 mm were made by pressing (at a

pressure of about 41.4 kPa for 30 seconds using a WABASHTM heated press, Wabash Co., Wabash, IN) about 10 g of the graft copolymers between chrome-coated aluminum plates at about 200°C. Pressed samples were 5 quenched from the molten state to the solid state in a room temperature water bath. Hexadecane contact angle measurements were made on these films and on polypropylene base polymer control sample using the sessile drop method as described by A. W. Newman and R. J. Good in "Techniques for Measuring Contact Angles,"

Surface and Colloid Science, 11, Plenum Press, NY, 1979. Results are shown in TABLE 2.

TABLE 2

15	Example	Contact Angle (°)
	PP Control	0
20	PTFE 1 2	47 ± 1 22 ± 3 19 ± 3
	3 4 5	20 ± 2 23 ± 3 21 ± 3
25	6 7	15 ± 2 20 ± 3
	8	19 ± 4 20 ± 2

The results in TABLE 2 show improved surface 30 activity, compared to a polypropylene (PP) control, for all Examples.

Electron spectroscopy for chemical analysis (ESCA) measurements were also done to confirm grafting. Fluorine to carbon (F:C) ratios for Examples 1 and 4 are listed in TABLE 3 below.

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TABLE 3

. *	Example	<u>c</u>	omposi	<u>tion</u>	F:C
	Control		PP		0
	1	*	PP/BuF	OSEA	0.25
	4		PP/BuF	OSEA	0.31

TABLE 4 below compares surface energy values determined for films of polypropylene, polytetrafluoro-ethylene (PTFE), and the graft copolymer of Example 2.

TABLE 4

Polymer Sample Surface Energy (Dyner Care) Polypropylene Control 32	<u>1)-</u>
Dalamananti ANA ('ARTEAL	
the many polypropylene concret, the property of the person of	
PTFE 22	- ''
5 Example 2	

The data in TABLE 4 indicate that the graft copolymer of Example 2 has a surface energy lower than the polypropylene control and approaching that of PTFE.

180° Peel Adhesion Test:

A 2.5 cm wide, 20.3 cm long strip of pressure-sensitive adhesive tape is adhered to a 10.1 cm wide, 15.2 cm
25 long sheet of a test substrate (a pressed sample of a graft copolymer of the invention) with a free end of the tape extending beyond the end of the test substrate. The tape/substrate sample is rolled twice with a 1.35 kg hard rubber roller to ensure contact
30 between the adhesive and the test substrate. The sample is aged at room temperature for 24 hours. The free end of the tape is removed from the test substrate by pulling at 180° at a rate of 15.2 cm/minute using a Slip/Peel Tester, (available from Instrumentors, Inc.,

180° Peel adhesion tests were run on pressed film samples of several examples and on an ungrafted polypropyl ne base polymer control sample using SCOTCHTM adhesive tape #8411 (3M) (an acrylate-based

pressure-sensitive adhesive with a 180° peel adhesion to glass of 110 g/cm) and 3M adhesive tape STA-115 (a KRATONTM rubber-based pressure-sensitive adhesive tape with a 180° peel adhesion to glass of 430 g/cm).

5 Results are shown in TABLE 5 below wherein each number represents the average of five independent determinations.

10	TAE	LE 5	
	Peel Forc		Peel Force
<u>Example</u>	(acrylate-ba	sed)	(KRATON TM-based)
<u> </u>	(g/cm)		(g/cm) 517
Control PP	209		311
15 1	198		. 330
	143		308

154

TABLE 5 indicates improvement in release
20 properties of graft copolymers of the invention
compared to the polypropylene control. Graft
copolymers of Examples 1, 2, and 3 show 5%, 32%, and
26% improvement in release properties for the
acrylate-based adhesive tape, respectively, and 36%,
25 40%, and 47% improvement in release properties for the
KRATONTM rubber-based adhesive tape.

A pressed film of the graft copolymer of
Example 1 was prepared as described above by pressing
between PTFE-coated aluminum plates at 200°C. The film
30 was quenched in a room temperature water bath. Samples
of the film were then annealed in an oven at 150°C for
various times. Hexadecane contact angle and ESCA
measurements were performed as described above.
Results are shown in TABLE 6 below.

TABLE 6

Annealing Time (minutes)	 adecan	The state of the s	F:C
0	 22 ± 1		0.25
5.0.5	22 ± 2		0.27
1.0	26 ± 3		0.24
2.5	32 ± 3		0.39
3.0	42 ± 4		
5.0	40 ± 4		
10 6.0	45 ± 3		0.45
8.0	47 ± 2		
23.0	46 ± 2		

The data in TABLE 6 demonstrate that

15 annealing the graft copolymers of the invention at

150°C enhances surface activity by increasing the
amount of fluorochemical at the surface of the graft
copolymer. The data above indicate an increase in
fluorochemical surface activity with increased

20 annealing time over about 6 minutes, after which time
there appears to be little advantage gained by further
annealing.

Example 10

N-Butyl perfluorooctanesulfonamidoethyl acrylate (BuFOSEA) was grafted to polypropylene (Himont PROFAXTM PP PF-301) as described in Example 1 above.

The fluorochemical olefin was added to the extruder at a rate of 2 mL/min (3.16 g/min) with a melt temperature of 180°C. Total flow rate was held at about 40 g/min.

The hexadecane contact angle of the resultant graft polymer was measured and found to be 39°. ESCA showed a F:C ratio of 0.20 on a pressed film prepared as described in Example 1 using PTFE-coated 35 aluminum plates.

Resistance to various solvents was also measured to determine stability of the surface to chemicals. 0.6 g - 0.8 g samples of the pressed film were placed in the solutions listed in TABLE 7 below.

40 After storage as indicated in TABLE 7, the film samples

were rins d and dried, and the percent weight loss was determined. Hexadecane c ntact angles and F:C ratios were measured for each film sample.

TABLE 7
CHEMICAL RESISTANCE OF FLUOROCHEMICAL GRAFT COPOLYMER

	Treatment Hexadecane (days) Wt. Loss (%) Contact Angle(°)	F:C Ratio
10	Polypropylene Control - 0	
	Control ⁴ 0% 32 ± 1	0.29
	20% HCl (5) 3.9% 43 ± 4	0.33
	25%H ₂ SO ₄ (5) 1.5% 42 ± 4	0.23
	2% Chromic 1.2% 49 ± 1	0.32
15	Acid (5)	
	3% NaOH (5) 0.1% 35 ± 1	0.23
1	Mineral Oil (5) 5.4% 24 ± 3	0.34
	Water (9) 0.1% 41 ± 3	0.31
	Air (9) 28 ± 3	0.45
20		0.45

A Treatment at room temperature. All others are at 82°C.

The data in TABLE 7 show that exposure to aqueous treatment decreases hexadecane contact angle, while treatment with hydrophobic organic materials increases hexadecane contact angle. However, in all cases, hexadecane contact angle remains between 20° and 50° after treatment. This indicates that oil repellency is maintained, and that the graft copolymers of the invention are resistant to chemical degradation under the indicated conditions.

35 Examples 11-46

Polypropylene/BuFOSEA graft copolymers were prepared as described in Examples 1-9, above except that in Examples 11-22 only LUPERSOLTM 101 initiator was used in the amounts set forth in TABLE 9; in Exampl s

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23-34 only LUPERSOLTM 130 initiator was used in the amounts set forth in TABLE 9; in Examples 35-46 a 1:1 mixture of LUPERSOLTM 101 initiator and LUPERSOLTM 130 initiator was used in the amounts set forth in TABLE 9. 5 Screw speed was 100 rpm, and monomer flow was 2 mL/min. The percent of grafted fluorochemical olefin was determined using x-ray fluorescence spectrometry to determine percent fluorine. Processing conditions, i.e., melt temperature and total flow, mole percent 10 BuFOSEA reacted, and weight percent grafted BuFOSEA in

the graft copolymers, are shown in TABLE 8.

			TABLE	8		
	. 0 <u> </u>	Initiator (Conc.	Melt Temp	Total Flow (q/min)	Mol % BuFOSEA Reacted	Wt % BuFOSE
5	Example 11	<pre>vt. %) L101(0.1)</pre>	(°C) 180	40.9	81	6.3
	12	u ,	200	348	70	6.4
	13	11	220	36.7	. 79	6.8
	14	L101(0.25)	180	30.3	78	8.1
	15	0	200	40.8	89	6.9
10	16	u 🛒	220	39.3	86	6.9
	17	L101(0.5)	180	40.6	91	7.1
	18		200	42.9	88	6.5
	19	10	220	46.3	86	5.9
	20	L101(1.0)	180	45.6	92	6.4
15	21		200	38.6	90	7.2
	22	•	220	53.6	95	8.9
	23	L130(0.1)	180	42.8	88	6.5
	24	er (200	46.7	96	6.5
	25		220	45.9	100	6.9
20	26	L130(0.25)	180	41.4	93.	7.1
ings Jagor	27	1	200	41.6	86	6.5
	28		220	47.6	88	5.8
	29	L130(0.5)	180	29.9	75	7.9
	30		200	43.4	93	6.8
25	31		220	43.3	99	7.2
	32	L130(1.0)	180	39.5	98	7.8
	33		200	24.7	70	9.0
	34		220	43.1	86	6.3
	35	L101/L130(0.1)	180	36.3	63	5.5
30	36		200	42.3	73	5.5
	37	er e	220	42.0	76	5.7
1	38	L101/L130(0.25	5) 180	34.2	64	5.9
	39		200	49.9	94	5.9
	40		220	37.5	73	6.1
35	41	L101/L130(0.50) 180	35.8	78	6.9
	42	11	200	-33-0	- 78-	7. -6
	43	. 1944 - 1944 - 1944 - 1944 - 1944 - 1944 - 1944 - 1944 - 1944 - 1944 - 1944 - 1944 - 1944 - 1944 - 1944 - 194	220	35.1	74	6.7
	44	L101/L130(1.0)	180	27.0	73	8.5
. (*	45	n ,	200	37.7	100	8.4
40	46	u ,	220	41.3	100	9.6
	Contro	1 -	180	40	0	0

TABLE 8 indicates that graft copolymers of this invention can be made at a wide variety of initiator concentrations. All Examples contained at least 5.0% (by weight) BuFOSEA, and grafting efficiency 5 was good in all Examples, ranging from 63% to 100%.

Example 47

FIG. 3 shows a twin screw extruder much like that shown in FIG. 2 and described in Examples 1-9

10 above. Particularly, the extruder comprises a feed zone 40, a heated barrel that comprises barrel section 42 comprising both an initiation/melt zone and a monomer addition zone, barrel sections 44, 46, and 48 comprising a reaction zone, barrel section 50

15 comprising a devolatilization zone and barrel section 52 comprising a block zone, and a die 54. Transducer ports (e.g., T₁ represents transducer number 1 in barrel section 42) are located as shown in FIG. 3, and thermocouple ports are located in each heated section of the extruder. The extruder was a LEISTRITZTM Model ASF67GG.

Temperature profile of the extruder was as shown in TABLE 9 below:

25 <u>TABLE 9</u>	
Transducer	T(°C)
	182
사람 이 자연의 시간 10 kg (1 kg - 2 kg) 시간 (1 kg - 2 kg) 시간 (1 kg - 2 kg) 시간 (1 kg - 2 kg - 2 kg) 시간 (1 kg - 2	171
	181
30 ·	179
	188
	181
	181
[2] [1] [1] [2] [2] [2] [2] [3] [3] [4] [4] [4] [4] [4] [4] [4] [4] [4] [4	180
35.	180
	194
	194

The feed zone was ambient temperature, and screw speed was 50 rpm.

Base polymer feed hopper and extruder feed throat were purged with nitrogen. Base polymer (DYPROTM 8771) was fed at a rate of 18.1 kg/h using a K-tron 6300 feeder. The initiator, a 1:1 mixture by weight of LUPERSOLTM 130 initiator and LUPERSOLTM 101 initiator, was purged with nitrogen and fed at a rate of 2 mL/min using a single piston RUSKATM positive displacement pump 10 at the downstream end of the feed throat, a distance of 270 mm from the start of the screws. BuFOSEA was purged with nitrogen and fed at a rate of 463 mL/h through a high pressure injection valve using a dual piston RUSKATM positive displacement pump, at a distance 15 of 610 mm from the start of the screws. Vacuum venting of unreacted BuFOSEA was performed in heated barrel section 50. Product graft copolymer was extruded through a 10-strand die that fed into a water bath and a CONAIRTM pelletizer.

20

Examples 48-56

Films of the graft copolymer were made as in Example 1 using chrome-coated aluminum plates at 190°C and a pressure of 837 mPA for 30 seconds. Also, films 25 were made by pressing the graft copolymer of this invention between plates that had been covered with polyimide tape (KAPTONTM Film Tape #5413, 3M) and plates that had been covered with PTFE tape (SCOTCHTM PTFE Film Tape #5490, 3M). Films of graft copolymers of Examples 1 and 4 were also made by pressing the graft copolymer of the invention between the chrome, polyimide, and PTFE surfaces as described above. Hexadecane contact angles were measured as described in Example 1 above. Results are shown in TABLE 10 wherein each result is an average of ten independent determinations.

TABLE 10

	<u>Example</u>	Polymer Sample PP Control	Hexadecan <u>Chrom</u> 0	e Contact e Polyi	Angle (°) mide PTFE O O
5	48	Example 1	16 <u>+</u>	3	
	49	Example 1		33	<u>+</u> 2
	50	Example 1			49 ± 5
	51	Example 4	13 ±	2	
10	52	Example 4		29	± .3
	53	Example 4			43 ± 4
	54 55	Example 47 Example 47	19 ±	2 27	± 1
15	56	Example 47			50 <u>+</u> 5

The results in TABLE 10 show that the graft copolymer of Example 47 exhibits surface activity similar to that 20 of the graft copolymers of Examples 1 and Example 4. The results in TABLE 10 also show the effect of the surface against which the graft copolymer is formed. The samples melted against the PTFE surface show higher surface activity than those formed against the more 25 polar surfaces polyimide and chrome.

Examples 57-59

The following examples describe graft polymerization of N-ethyl

perfluoroctanesulfonamidoethyl acrylate (EtFOSEA) to polypropylene. Polypropylene (Amoco PP 5219, Amoco Chemical Naperville, IL) was premixed with 0.25 wt % LUPERSOLTM 101 initiator, 0.25 wt% LUPERSOLTM 130 initiator, and 5.0 wt% EtFOSEA. The mixture was purged with nitrogen, added in the feed zone, extruded, and collected as described in Example 1. Screw speed of the extruder was 100 rpm. TABLE 11 lists the process conditions for Examples 57-59.

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TABLE 11

		Total Flow Rate
<u>Example</u>	Melt Temp. (°C)	<u>(g/min)</u>
	180	40.6
	선생님들은 어느로 들어서 되는 것이 바다	
5 58	200	43.7
50	220	40.5

Examples 60-65

Films of the graft copolymers of Examples

10 57-59 were made as described in Example 1 by pressing
between chrome surfaces at 185°C and 2790 mPA for 60
seconds, and by pressing between PTFE surfaces at 174°C
and 1046 mPA for 30 seconds as indicated in TABLE 12
below. Hexadecane contact angles were measured and are
15 shown in TABLE 12, wherein each entry represents the
average of ten independent determinations.

TABLE 12

20 <u>Example</u>	Polymer Hexadecane Contact Angle (°) Sample Chrome PTFE	
	PP Control 0	
60	Example 57 25 ± 2	
61	Example 57 65 ± 2	
		4
25 62	Example 58 24 ± 3	
63	Example 58	
64	Example 59 31 ± 1	_
65	Example 59 71 ± 3	

The data in TABLE 12 indicate that the pressed films of the invention have high surface activity compared to that of the polypropylene control.

Also, the PTFE-pressed films have a twofold or greater hexadecane contact angle than the chrome-pressed films.

Examples 66-67

30

The following Examples describe the graft polymerization of N-ethyl

perfluorooctanesulfonamidoethyl methacrylate (EtFOSEMA) to polypropylene.

Polypropylene (Amoco PP 5219, Amoco Chemical, Naperville, IL)/EtFOSEMA graft copolymers

were prepared as described in Examples 1-9 above. The EtFOSEMA was used as a 1:1 solution by weight in tetrahydrofuran, and the solution was fed at 4 mL/min. Melt temperatures were 180°C for Example 66 and 200°C for Example 67. Tetrahydrofuran and unreacted monomer were vented as in Examples 1-9. Screw speed was 60 rpm. Polymer flow rates were 38.3 (Example 66) and 42.4 (Example 67) g/min.

Examples 68-71

of films prepared from the graft copolymers of Examples 66 and 67. Results indicate improved surface activity compared to the control sample, and a higher surface activity for the PTFE-pressed films compared to the cohome-pressed films.

TABLE 13

<u>Example</u>	Polymer Sample	Hexadecane <u>Chrome</u>	Contact	Angle (°) PTFE
25	PP Contro	o1		0
68	Example	66 14 <u>+</u> 2		
69	Example (66		24 ± 3
70	Example	67 14 ± 3		
30 71	Example	67		30 ± 3

Examples 72-73

The following examples describe graft polymerization of N-allyl

perfluoroctanesulfonamidomethyl ethane (AlfOSME) to polypropylene (Amoco PP 5219).

These graft copolymers were prepared as in Examples 66 and 67 with melt temperatures of 180°C

(Example 72) and 200°C (Example 73) and polymer flow rates of 41.7 (Example 72) and 42.3 (Example 73) g/min.

Examples 74-77

5 TABLE 14 lists hexadecane contact angles on films of the graft copolymers of Examples 72 and 73, prepared as in Examples 60-65. Again, increased fluorochemical surface activity compared to the control sample was seen, and the PTFE-pressed films showed 10 higher surface activity than the chrome-pressed films.

TABLE 14

<u>Example</u>	Polymer Sample	Hexadecane <u>Chrome</u>	Contact	Angle (°)
15	PP Control	0	N (X)	0
74	Example 72	15 ± 1		
75	Example 72			45 ± 3
76	Example 73	22 ± 3		
20 77	Example 73			42 ± 3

Examples 78-83

The following Examples describe the preparation of fluorochemical graft copolymers using low molecular weight base polymers. The graft copolymers were prepared according to the general method of Examples 1-9 above using a 1:1 mixture of LUPERSOLTM 101 initiator and LUPERSOLTM 130 initiator as the initiator system and the components and conditions listed in TABLE 15 below. The fluorochemical olefins were used in an amount of 10 percent by weight based on the weight of the base polymer. Initiator concentration is based on the weight of the base polymer.

The graft copolymers were extruded into film substrates of 0.1 mm thickness. ESCA analysis and 180° peel adhesion results (using the test described above in connection with TABLE 5 and 3M Box Sealing Tape #371, a KRATONTM rubber-based adhesive tape with a peel

strength of 610 g/cm when adhered to glass) are shown in TABLE 15.

The ESCA data of TABLE 15 indicate that there is more fluorine on the surface of the films than would be expected based on the amount of fluorochemical olefin used in preparing the graft copolymer. The release data indicate in all cases that the initial peel force of the films was superior to that of control base polymer.

CABLE 15

Example	Base Polymer (MFI ^c)	Fluorochemical Olefin	Initiator Concentration (%)	ESCA (%F)	Peel Force (q/cm) Initial Aced ^D	e (g/cm)
PP control	Exxon ^A 3085 (35)			0	500	550
78	Exxon ^A 3085 (35)	Etfosea	.	39	8	560
79	Exxon ^A 3085 (35)	Etfosea	0.5	47	310	460
08	Exxon ^A 3085 (35)	Etfosema		33	140	630
PE control	Dow ⁸ 40060M (40)			0	160	510
81	Dow ^B 40060M (40)	Etfosea	0.25	38	20	250
82	Dow ^B 40060M (40)	Etfosea	0.12	22	20	170
83	Dow ^B 40060M (40)	Etfosema	0.25	48	190	340

Exxon Chemical Co., Houston, TX

Dow Chemical Co., Midland, MI

Melt flow index

"Aged" peel force was determined using the 180° Peel Test after heating the tape/substrate sample at 43°C for 11 days.

Examples 78-80 do not have particularly durable aged peel force. In the case of Examples 81-83, however, aged peel force was significantly lower than that of the control polyethylene base polymer and in the range suitable for commercial tape applications.

Examples 84-88

Polymer blend films were made by dry-blending 10 parts by weight of a graft copolymer of 10 the invention with 90 parts by weight of a pure matrix polymer, adding the resulting mixture to the hopper of a 1 1/4 inch (5 cm) Killion extruder (Killion Extruders, Inc., Verona, NJ), and extruding into a 0.1 mm thick, 15.2 cm wide film substrate under the 15 following conditions:

Extruder temperatures:	Zone 1 149°C
	Zone 2 201°C
	Zone 3 220°C
	Zone 4 221°C
	Neck tube 218°C
	Die 218°C

Extruder rpm: 25
25 Casting roll temperature: 15°C

Film substrates were made using the components set forth in TABLE 16 below. ESCA results and 180° peel adhesion results are also shown in TABLE 16.

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<u>Example</u>		Graft Copolymer	ESCA (%F) FO	ged Peel erce (g/cm) ^c
5 PE control	Dow ^A 6806		0 .	540
84	Dow ^A 6806	Example	81 15	400
PP control	Fina ^B 3374X		0	580
85	Fina ^B 3374X	Example	83 17	260
LO 86	Fina ^B 3374X	Example	79 23	340
87	Fina ^B 3374X	Example	78 18	220
88	Fina ^B 3374X	Example	82 14	260

- A. Dow Chemical Co., Midland, MI
- 15 B. Fina Chemical Co.
 - C. "Aged" peel force was determined using the 180°
 Peel Test set forth in Examples 1-9 above (3M
 adhesive tape STA-115) after heating the
 tape/substrate sample at 43°C for 11 days.

20

The ESCA data of TABLE 16 indicate that there is more fluorine on the surface of the films than would be expected based on the amount of graft copolymer used in preparing the blend. The release data indicate in all cases the aged peel force of the blend is lower than that of the control base polymer and in the range suitable for commercial tape applications.

Various modifications and alterations of this invention will become apparent to those skilled in the art without departing from the scope and spirit of this invention, and it should be understood that this invention is not to be unduly limited to the embodiments set forth herein.

30

35

The Claimed Invention Is:

- 1. A fluorochemical graft copolymer comprising: a base polymer comprising polymerized units derived from monomers having terminal olefinic double bonds, having a moiety comprising a fluoroaliphatic group grafted thereto.
- A graft copolymer according to Claim 1,
 wherein the grafted moiety is a fluorochemical olefin comprising a fluoroaliphatic group and a polymerizable double bond.
- 3. A graft copolymer according to Claim 1,
 15 wherein the base polymer is selected from the group
 consisting of polymethyl methacrylate, poly4-methylpentene, polypropylene, polybutylene,
 polystyrene, polyethylene, polybutadiene,
 ethylene/vinyl acetate copolymer, ethylene/butyl
 20 acrylate copolymer, and mixtures and blends thereof.
 - 4. A graft copolymer according to Claim 1, wherein the grafted moiety is a fluorochemical olefin of the formula

 $(R_f)_aQ(CR=CH_2)_b$

wherein R is hydrogen, trifluoromethyl, or lower alkyl;

- a is an integer from 1 to about 10;
- b is an integer from 1 to about 6;
- Q is an (a+b)-valent linking group that does not substantially interfere with free radical polymerization; and
- R_f is a fluoroaliphatic group comprising a fully fluorinated terminal group containing at least seven fluorine atoms.

- 5. A graft copolymer according to Claim 4, wherein the grafted moiety is a fluorochemical olefin of the formula $C_8F_{17}SO_2N(C_4H_9)CH_2CH_2OC(0)CH=CH_2$.
- 6. A graft copolymer according to Claim 4, wherein the grafted moiety is a fluorochemical olefin of the formula $C_8F_{17}SO_2N(C_2H_5)CH_2CH_2OC(O)CH=CH_2$.
- 7. A graft copolymer according to Claim 4, 10 wherein the grafted moiety is a fluorochemical olefin of the formula $C_8F_{17}SO_2N(C_2H_5)CH_2CH=CH_2$.
 - 8. A graft copolymer according to Claim 4, wherein the grafted moiety is a fluorochemical olefin of the formula $C_8F_{17}SO_2N(C_2H_5)CH_2CH_2OC(O)C(CH_3)=CH_2$.
 - 9. A graft copolymer according to Claim 1, wherein the base polymer is a low molecular weight polyolefin.
 - 10. A graft copolymer according to Claim 9, wherein the base polymer has a melt index of at least about 20.
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 11. A fluorochemical graft copolymer
 according to Claim 1 comprising a polymeric backbone
 comprising polymerized units derived from monomers
 having terminal olefinic double bonds, having grafted
 thereto a moiety of the formula

30 -- CH2-CHR-Q-Rf

wherein R is hydrogen, trifluoromethyl, or lower alkyl, Q is a divalent organic linking group that does not interfere with free-radical polymerization, and R_f is a fluoroaliphatic group comprising a fully fluorinated terminal group containing at least seven fluorine atoms.

- 12. A graft copolymer according to Claim 1 in the form of a film or a coating.
- 13. A process for preparing a fluorochemical graft copolymer according to Claim 1, which process comprises:
 - (1) feeding to a reactor materials comprising:

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- (a) a base polymer comprising polymerized units derived from monomers having terminal olefinic double bonds;
- (b) an effective amount of a free radical initiator system comprising one or more free radical initiators; and
- (c) a fluorochemical olefin comprising a fluoroaliphatic group and a polymerizable double bond, wherein all materials are substantially free of oxygen;
- (2) reacting the materials in the reactor to provide the fluorochemical graft copolymer; and
- (3) withdrawing the fluorochemical graft copolymer from the reactor.
 - 14. An annealing method for increasing the amount of grafted fluorochemical at the surface of a composition comprising a fluorochemical graft copolymer, comprising the steps of:
 - a) providing a surface comprising a fluorochemical graft copolymer according to Claim 1, and
- b) annealing the surface by heating it at a 35 temperature and for a time effective to increase the amount of the grafted fluorochemical at the surface.
 - 15. A forming method for controlling the amount of grafted fluorochemical at the surface of a

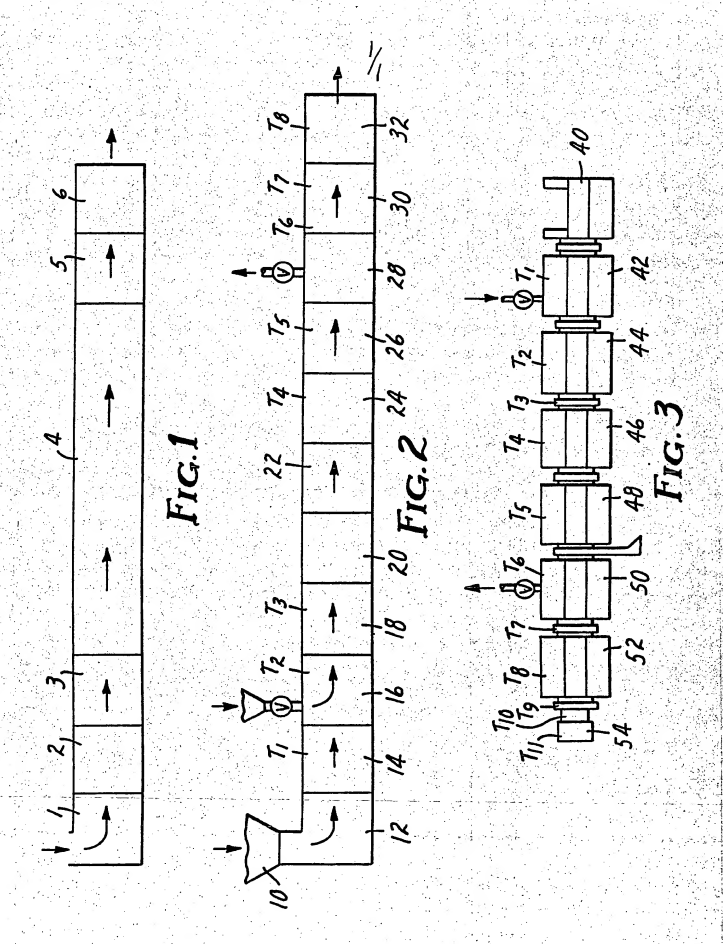
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composition comprising a graft copolymer according to Claim 1, comprising the steps of:

- (1) selecting a surface that is made of a material that will control the amount of grafted fluorochemical at the surface of the composition; and
- (2) forming the composition against the surface selected in step (1).
- 16. A polymer blend comprising a fluorochemical graft copolymer according to Claim 1 and a matrix polymer that is miscible with the base polymer of the graft copolymer.
- 17. A polymer blend according to Claim 16, wherein the base polymer is a low molecular weight polyolefin.
 - 18. A polymer blend according to Claim 17, wherein the base polymer has a melt index of at least about 20.
 - 19. A polymer blend according to Claim 16 comprising at least about 1 percent of the graft copolymer by weight based on the weight of the matrix polymer.
- 20. A polymer blend according to Claim 16 comprising at least about 10 percent of the graft copolymer by weight based on the weight of the matrix polymer.
 - 21. A polymer blend according to Claim 16 comprising at least about 30 percent of the graft copolymer by weight based on the weight of the matrix polymer.
 - 22. A polymer blend according to Claim 16 in the form of a film or a coating.

23. A method f r reducing the surface energy of a polymer comprising polymerized units derived from monomers having terminal olefinic double bonds, comprising the step of grafting to said polymer a fluorochemical olefin comprising a fluoroaliphatic group and a polymerizable double bond.

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	INTERNATIONAL SEARCH REPORT	US 92/00591
		03 32,00032
I. CLASSIFICATI	N F SUBJECT MATTER (if several classification symbols apply, Indicate all) 6	
IPC5: C 08 F	national Patent Classification (IPC) or to both National Classification and IPC 255/00	
II. FIELDS SEARC	CHED	
	Minimum Documentation Searched ⁷	
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1, 63	Documentation Searched other than Minimum Documentation to the Extent that such Documents are Included in Fields Searched ⁸	
	CONSIDERED TO BE RELEVANT ⁹	
Category Ci	ation of Document, ¹¹ with indication, where appropriate, of the relevant passages ¹²	Relevant to Claim No.13
X US,	A, 4666991 (MATSUI ET AL) 19 May 1987, see column 3, line 3 - line 23; column 5, line 63 - column 6, line 7; claims 1-4,7-8	1-4,9- 13,16, 23
	A, 4590236 (KÖNIG ET AL) 20 May 1986, see column 2, line 17 - line 39; column 2, line 62 - column 3, line 42; claims 1,4	1-13,16, 23
x Us,	A, 4100225 (MUELLER) 11 July 1978, see column 4, line 57 - column 5, line 22; column 6, line 42 - column 7, line 17; claims 1-12	1-13,16, 23
"A" document considered "E" earlier doc filing date "L" document which is citation or "O" document other mea "P" document later than IV. CERTIFICAT Date of the Actual 10th June International Sear	which may throw doubts on priority claim(s) or led to establish the publication date of another other special reason (as specified) referring to an oral disclosure, use, exhibition or not published prior to the international filing date but the priority date claimed ON Completion of the International Search Date of Mailing of this International	nce, the claimed invention cannot be considered to nce, the claimed invention we an inventive step when the ne or more other such docup obvious to a person skilled ne patent family

International Application No. PCT/US 92/00591

II. DOCI	MENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)	* .
Category *	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to Claim No
X	US, A, 3876729 (MUELLER) 8 April 1975, see column 1, line 10 - line 20; claim 1	1-13,16, 23
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ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.PCT/US 92/00591

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This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 30/04/92.

The European Patent office is in no way liable for these particulars which are merely given for the purpose of information.

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